Direct observation of processive exoribonuclease motion using optical tweezers

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Bacterial RNases catalyze the turnover of RNA and are essential for gene expression and quality surveillance of transcripts. In Escherichia coli, the exoribonucleases RNase R and polynucleotide phosphorylase (PNPase) play critical roles in degrading RNA. Here, we developed an optical-trapping assay to monitor the translocation of individual enzymes along RNA-based substrates. Single-molecule records of motion reveal RNase R to be highly processive: one molecule can unwind over 500 bp of a structured substrate. However, enzyme progress is interrupted by pausing and stalling events that can slow degradation in a sequence-dependent fashion. We found that the distance traveled by PNPase through structured RNA is dependent on the A-U content of the substrate, and the removal of its KH and S1 RNA-binding domains can reduce enzyme processivity without affecting the velocity. By a periodogram analysis of single-molecule records, we establish that PNPase takes discrete steps of six or seven nucleotides. These findings, in combination with previous structural and biochemical data, support an asymmetric inchworm mechanism for PNPase motion. The assay developed here for RNase R and PNPase is well suited to studies of other exonucleases and helicases.

Significance

Bacteria regulate the synthesis and degradation of RNA molecules to ensure timely and robust responses to an ever-changing environment. The transcript’s lifetime can be influenced profoundly by a secondary structure that can form in the RNA and that may inhibit or promote its digestion by RNases. The molecular mechanisms by which RNases interact with structured RNAs therefore are of great interest. In this study we used optical tweezers to investigate the mechanistic properties of two such enzymes from Escherichia coli, polynucleotide phosphorylase and RNase R. Our results offer new insights into the functional characteristics of these two enzymes, including the sequence-dependent behavior of RNase R and the presence of discrete steps of six or seven nucleotides taken by polynucleotide phosphorylase.
more limited processivity, with an individual enzyme capable of unwinding around 60 bp of structured RNA, provided that it is rich in adenine and uracil. PNPase mutants lacking the KH and S1 domains exhibited reduced processivity but moved along dsRNA at rates similar to the WT enzyme. Finally, we find that PNPase translocates in discrete steps of six or seven nucleotides, a behavior that suggests it may advance by an asymmetric inchworm mechanism. The single-molecule assay developed for this work should be readily adaptable to the study of additional exonucleases and helicases.

**Results**

In the exoribonuclease assay, a substrate RNA is hybridized to a complementary strand composed of either DNA or RNA to form a duplex. At its 3′ end the duplex carries a short (12- to 14-nt) noncomplementary loading sequence, which overhangs and serves as a target for the initial binding of the enzyme. The hybridized duplex and associated loading sequence are suspended between two optically trapped polystyrene beads using chemical linkages and a DNA “handle” (Fig. 1). Enzyme-driven unwinding of the duplex region, driven by hydrolysis of the bound strand, converts the double-stranded substrate into a single-stranded form that has a longer contour length, resulting in a change in the end-to-end extension by an amount related to the applied load (−0.2 nm per nucleotide at 20 pN): an RNA duplex has a rise of 0.28 nm per nucleotide (32), whereas ssRNA (or ssDNA) has a force-dependent rise of 0.48 nm per nucleotide at 20 pN (33). Using this approach, we were able to monitor the activity ofRNase R and PNPase in real time and at near-nucleotide resolution.

**RNase R: Velocity, Processivity, and Pausing Behavior.** RNase R was assayed on a 155-bp dsRNA construct held at 20 pN with an optical force clamp. The motion was highly processive, with 77% of enzymes (n = 10 of 13) advancing through more than 24 nm and nearly half (46%) unwinding the entire substrate (Fig. 2A). Because the complete unwinding of the RNA substrate corresponded to an extension change of 26.9 ± 0.2 nm (mean ± SE), we calculate that the release of each nucleotide increased the tether length by 0.174 ± 0.001 nm (mean ± SE). This value is close to the predicted value of 0.2 nm per nucleotide (32, 33), and it was used to calculate the number of nucleotides unwound in subsequent experiments with dsRNA. RNase R was similarly processive when degrading RNA hybridized to a DNA complement, with 75% of molecules (n = 11 of 15) advancing over 24 nm, and 45% removing the entire transcript. The conversion factor for unwinding a DNA–RNA hybrid duplex was 0.178 ± 0.004 nm per nucleotide (mean ± SE), the same, within error, as for dsDNA.

RNase R moved along dsRNA with an average velocity of 88 ± 8 nt/s (mean ± SE), but its motion was not uniform: Most enzymes paused briefly at one or more locations, a behavior reminiscent of elongation by RNA polymerase (34). However, the pauses observed were confined primarily to the first ~16 nm of the dsRNA substrate, and pauses with durations greater than 0.1 s were not found in the ~9-nm (50-nt) stretch immediately following this region. Based on the rates of traversal of this comparatively pause-free region, we estimate the pause-free velocity of RNase R to be 151 ± 11 nt/s (SI Materials and Methods). In contrast, the average velocity of RNase R on the DNA–RNA hybrid was significantly faster, 128 ± 14 nt/s (two-tailed t test, P = 0.016), and the corresponding pause-free velocity was 184 ± 20 nt/s (22% faster than on dsRNA). Because RNase R digests an identical RNA sequence in both experiments, the slower velocity over dsRNA must be a consequence of the enzyme’s being differentially sensitive to the structure (35–37) or of the stability (38) of the DNA–RNA hybrid.

RNase R paused with high probability at three specific positions on the dsRNA construct (Fig. 2B and Fig. S1). By aligning single-molecule records, we determined that pauses occurred at extensions of 6.0 ± 0.1 nm (34–36 nt), 12.4 ± 0.1 nm (70–74 nt), and 15.4 ± 0.1 nm (87–91 nt) relative to the starting position. There were no obvious sequence similarities among these three locations, nor did we observe pausing behavior at the equivalent sequence positions on the DNA–RNA hybrid. To confirm that the pauses on dsRNA were sequence dependent, we repeated the experiment on a 1,500-bp dsRNA substrate carrying three tandem repeats of the original 155-bp sequence, each separated by a spacer region of six base pairs (Fig. 2C and Fig. S2). For the enzymes that successfully traversed all three tandem repeats (n = 7 of 18), pauses were observed at each of the expected positions in every segment (Fig. 2D).

The pauses scored in the tandem repeats enabled us to obtain an independent estimate of the conversion factor for relating the change in distance (measured in nanometers) to the number of nucleotides unwound for dsRNA. Our original estimate of 0.174 ± 0.001 nm per nucleotide (mean ± SE) had been calculated based on an assumption that RNase R completely digested the target substrate for the 155-bp construct. By instead measuring the change in distance between the corresponding pauses in tandem repeats, we independently calculated a value of 0.172 ± 0.002 nm per nucleotide (mean ± SE), which is identical within error.

With some probability, the motion of RNase R arrested completely at pause sites; the presence of multiple pause sequences engineered into the 1,500-bp construct likely hindered its full unwinding. Nonetheless, about one third of the enzymes successfully traversed at least 800 bp of dsRNA (n = 5 of 18). We therefore decided to examine RNase R behavior on a second 1,500-bp construct lacking the tandem pause elements to obtain an independent estimate of processivity. On this substrate, RNase R unwound an average of 503 ± 110 bp (mean ± SE; n = 16).

**PNPase: Stepping Behavior and Sequence Dependence.** We also studied the behavior of PNPase as it degraded the same (155-bp) dsRNA and DNA–RNA hybrid sequences used for the RNase R study and under an identical 20 pN load. These experiments were performed under buffer conditions that favored phosphorolysis (Materials and Methods). Under these conditions, PNPase removed an average of 23 ± 2 nt (mean ± SE; n = 20) from the dsRNA substrate and the same number of nucleotides (23 ± 1 nt; mean ± SE; n = 14) from the DNA–RNA hybrid (Fig. 3D). The average velocity of PNPase over these two substrates was the same within error: 121 ± 30 nt/s (mean ± SE; n = 15) for dsRNA and 129 ± 40 nt/s (mean ± SE; n = 10) for DNA–RNA.

The 155-nt substrate had an A+U content of 46%, overall. However, this content was not uniformly distributed but was enriched to a concentration of 69% in the 26-bp region initially digested by PNPase. To investigate whether the processivity of PNPase depended on the base composition, we studied its motion
on a DNA–RNA hybrid enriched for A+U in which the first 48 bp were replaced with a randomized sequence so that the first 50 bp carried an A+U concentration of 70% (hereafter referred to as “AU50”). On AU50, PNPase moved an average of 62 ± 3 nt (mean ± SE; n = 27), unwinding more than twice as much RNA as before while moving at the same rate, within error (137 ± 8 nt/s; mean ± SE).

In single-molecule records of PNPase motion on AU50, we occasionally observed brief pauses that were highly suggestive of enzyme stepping. Although the signal-to-noise ratio of individual records was not sufficiently good for the direct detection of steps, we were able to perform a periodogram analysis of the combined datasets (i.e., a Fourier analysis of the distribution of extension changes) to investigate the frequency of the stepwise events (39). The analysis yielded a strong peak at a spatial frequency of 0.87 ± 0.14 nm⁻¹ (mean ± SD), corresponding to an underlying step size of 6.6 ± 1.1 nt (Fig. 3B). We separately analyzed PNPase records collected on the original 155-bp construct (Fig. 3C) and obtained a peak at 0.94 ± 0.22 nm⁻¹ (mean ± SD) corresponding to a step size of 6.1 ± 1.4 nt, which was the same, within error. We conclude that the observed periodicity is unlikely to be a consequence of any sequence-dependent pausing but instead reflects a series of discrete steps of six to seven nucleotides taken by PNPase molecules. As an additional control, we carried out an analogous analysis of records obtained with RNase R (Fig. S3). Although those data were collected and analyzed using the identical apparatus, software, and sampling rate (2 kHz) and with enzymes that moved at similar velocities, we saw no evidence for such steps of six or seven nucleotides in the records of RNase R.

PNPase mutants in which the KH and S1 RNA-binding domains were deleted have been reported to display reduced nucleolytic activity (17, 40). We considered whether removal of these domains were deleted have been reported to display reduced nucleolytic activity (17, 40). We considered whether removal of these domains would have any effect on velocity or processivity. Using a PNPase ΔKH–ΔS1 mutant, we observed a twofold reduction in processivity on dsRNA (Fig. S4) relative to WT (14 ± 1 bp; mean ± SE; n = 10), but the velocity was the same within error (137 ± 43 nt/s; mean ± SE). Our findings suggest that the ~100-fold loss of nuclease activity reported for truncated mutants (40) may be a consequence primarily of reduced binding affinity, because a twofold drop in processivity alone, with no concomitant change in velocity, is insufficient to explain the pronounced loss of activity displayed by the mutants. This explanation is consistent with the comparatively infrequent observation of records of unwinding, which require productive binding of the enzyme, in assays using these mutants, in which higher concentrations of the enzyme (roughly eightfold greater than wild-type) were routinely required. The reduced processivity of the PNPase mutant led to short records of motion that precluded a periodogram analysis.

Discussion

We developed a high-resolution single-molecule assay for studies of helicases and exonucleases and used it to examine the RNA-unwinding activities of RNase R and PNPase. The experimental

Fig. 2. RNase R pauses at specific sites as it unwinds dsRNA. (A) Six representative single-molecule records of RNase R activity on the 155-bp dsRNA construct. The legend gives the three main pause locations (mean ± SE). (B) Three representative single-molecule records of RNase R activity on a 1,500-bp dsRNA construct carrying three tandem repeats (TR1–TR3) of the 155-bp sequence (Fig. S2). Approximate beginning and ending locations for each repeat are indicated (dashed lines). (C) Positional histogram of aligned RNase R records (n = 18) for the 1,500-bp construct (blue curve), including data from molecules that did not fully traverse the region (Materials and Methods). The predicted locations of pauses within the repeats are indicated (dark gray bars); these records were left-aligned at the measured location of the first pause (light gray bar at 11.4 ± 0.3 nm), which was found within 1 nm of its predicted location, at 10.5 ± 0.1 nm. The widths of the gray bars (2.0 nm) correspond to the approximate widths of peaks in the pause distributions.

Fig. 3. PNPase motion along RNA. (A) Twenty-two representative single-molecule records of PNPase advancing on 155-bp DNA–RNA hybrid constructs. Enzymes traversing RNA with 70% A+U content over the first 26 nt (blue traces, n = 11) moved an average distance of 23 ± 1 nt (mean ± SE). Enzymes traversing RNA with 70% A+U content over the first 50 nt (AU50; red traces, n = 11) moved a longer average distance of 62 ± 3 nt (mean ± SE). Records were smoothed with a 100-point boxcar window (t = 50 ms). (B) Periodogram analysis of records from the AU50 construct (n = 27) yielded a peak centered at 0.87 ± 0.14 nm⁻¹ (mean ± SD), corresponding to a step size of 6.6 ± 1.1 nt. (C) Periodogram analysis of records of the unmodified 155-bp construct (n = 25; data from the dsRNA and DNA–RNA hybrid are combined) yielded a peak centered at 0.94 ± 0.22 nm⁻¹ (6.1 ± 1.4 nt; mean ± SD).
The load applied to RNA was well below that required for spontaneous unwinding as the result of force-induced shearing (>25 pN) (41). Therefore, given the absence of a separate source of energy, the processive unwinding of RNA must be driven by the degradation of the RNA substrate. Nevertheless, the application of an external load is expected to perturb the energy landscape for the unwinding reaction, favoring the melting of any double-stranded regions and thereby influencing the velocity and processivity of PNPase and RNase R. This effect may explain why we observed greater processivity for PNPase than suggested by a previous estimate (18). Because the external load primarily affects the melting of the duplex, RNase R and PNPase are expected to move at least as fast and as far on ssRNA where no additional energy is required for unwinding. Therefore, the velocity and processivity values returned by our assay likely represent lower bounds for single-stranded substrates and upper bounds for double-stranded substrates.

The results reveal RNase R to be highly processive. The enzyme moves fairly rapidly and at a rate sensitive to whether it is acting on dsRNA or a DNA–RNA hybrid. We found that the propensity to pause (or arrest) is sensitive to sequence, and we identified three pause sites within the coding region of the podd gene from E. coli. The existence of pauses suggests that some RNAs may carry sequence elements that render them comparatively less sensitive to degradation by RNase R and therefore may indicate a role for this enzyme in the maturation of transcripts.

One recent study using single-molecule FRET (24) reported that both RNase R and its eukaryotic homolog, Rrp44, take steps of approximately four nucleotides along RNA. However, a periodogram analysis of our RNase R records did not yield a step of this size (Fig. S3), although the same approach successfully identified steps of six or seven nucleotides for PNPase. The discrepancy in the behavior of RNase R might be attributable to the different reaction conditions; we studied RNase R at 26 °C, but the FRET study was performed at a comparatively low temperature (12 °C) at which the enzyme velocity was reported to be approximately 50-fold slower than in our assay (41).

For PNPase, we found that the velocity and processivity were independent of whether it unwound dsRNA or a DNA–RNA hybrid. However, processivity was strongly influenced by the A+U content of the substrate, a result consistent with previous reports that the susceptibility of RNA hairpins to PNPase digestion depends on their thermodynamic stability (20).

Immediately following the initial phase of unwinding by PNPase, we typically observed an ~2 nm decrease in records of the tether extension, which took place on a timescale of 0.1–0.2 s, resulting in an apparent backtrack motion (Fig. 3A). Because the PNPase catalytic site is buried deep inside the enzyme core and therefore is spatially distinct from the point where the protein initially binds RNA, tether shortening may occur when PNPase releases a short fragment of unwound RNA that subsequently rehybridizes to its complement. In support of this explanation, we note that the average backtrack distance corresponded to the release of 12 ± 1 nt (mean ± SE; n = 44), as is consistent with current models for the PNPase–RNA interaction ahead of the catalytic site (16). The observation of steps of six or seven nucleotides, combined with structural and biochemical data for PNPase, enables us to postulate a mechanism by which this protein may advance along RNA (Fig. 4). In this asymmetric inchworm model, the KH and/or S1 domains bind to RNA on the 5′ side of the catalytic core and (for structured substrates) melt six or seven base pairs. The catalytic domains in the enzyme core then degrade RNA on the 3′ side, one nucleotide at a time, advancing until they encounter the bound KH/S1 domains. The latter domains then dissociate from the transcript and rebind downstream in the 5′ direction, repeating the cycle. In this model, the increases in tether extension observed in our single-molecule records correspond to the abrupt melting steps of approximately six or seven nucleotides, and the pauses between steps correspond to the degradation of the RNA by the PNPase core domains. The model relies on the presence of flexible linkers connecting the KH and S1 domains to the remainder of the protein; structural data suggest that these linkers could provide the comparatively large conformational changes required to execute a step of six or seven nucleotides (16). The precise roles of the individual RNA-binding domains and whether they bind via a concerted (as depicted in Fig. 4) or a hand-over-hand mechanism (16) remain to be determined.

In summary, the single-molecule assay presented here is well suited for studying the nanomechanical behavior of helicases and exonucleases, including the measurement of such properties as velocity, processivity, step size, and sequence-dependent effects. The assay is simple and convenient, and it has the advantage that WT motors can be studied without any of the chemical modifications required for fluorescent labeling or direct surface attachments. We anticipate that this approach should be applicable to the study of other processive exoribonucleases at the single-molecule level, including the human and archaeal exosomes, RNase II and RNase J.

Materials and Methods

RNases. PNPase and the truncated core of PNPase lacking the S1 and KH domains were purified as described (42). Ribonuclease R (RNOR7250; Epicentre) was stored at −20 °C.

DNA/RNA Constructs. Sequences and oligomers are described in SI Materials and Methods. Except where otherwise described, RNA was obtained by catalytic core digests and releases individual nucleoside monophosphates as it advances 3′ to 5′ until it reaches the RNA-binding domains. When the core catches up, the KH/S1 domains are released and become available to rebind further downstream along the RNA to complete additional cycles of degradation.

Fig. 4. Asymmetric inchworm model for PNPase. (A) PNPase has three KH/S1 RNA-binding domains (green rods), each of which is loosely coupled to its catalytic core via flexible linkers (black). (B) These domains bind to the RNA substrate downstream of the catalytic site, where they melt six or seven base pairs of dsRNA (digested strand in orange, complementary strand in red). (C) The catalytic core degrades RNA in a 5′ to 3′ direction, advancing until it reaches the RNA-binding domains. When the core catches up, the KH/S1 domains are released and become available to rebind further downstream along the RNA to complete additional cycles of degradation.
in vitro transcription (MEGAscript T7 kit, AM1333M, Life Technologies) of a DNA template, derived by PCR, from a 155-bp segment of the pALB3 plasmid (44). For corresponding in vitro mRNA, the 1,500-bp dsRNA construct, a second
biotin tag (Integrated DNA Technologies) to a transcript synthesized in vitro. The full-length RNA was purified by PAGE.

Supporting Information


Supporting Information

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SI Materials and Methods

Substrate RNA. Except where otherwise noted, all PNPase and RNase R data were collected using an RNA substrate with the sequence 5′-GGCCUGACAUAGAGUUCCGGAACCGGUGGUUUGACUGGAGAAUGCAGAUAACUGGGCGGUGACCUCUUUCGGCCACCGGUGUUAAACUGCGUGUCUCUGUUAGAAUGACUGGAGUAAAGCAGGUGCCUGUAGCAUUUGCGGAGCUAGGCAGCGGAGAAGCACC-GUAAAGACAUUUAGACAGAGAAGAGAAAGA-3′.

This RNA was generated by in vitro transcription of a DNA template, which was obtained by PCR of the pALB3 plasmid using the primers RNase R forward: 5′-AAAAAGAGTATAATACGACCTACTATAGGCTGACTTCTTGT-TG-3′ and 19rep-motor reverse: 5′-ATGGCATTTATATACGACCTACTATAGGCTGACTTCTTGT-TG-3′.

Substrate Complement: DNA. For assays performed on DNA–RNA hybrids, the strand complementary to the enzyme substrate carried a 5′-biotin tag, and was chemically synthesized (Integrated DNA Technologies). The sequence was as follows (bases complementary to the substrate RNA are underscored): 5′-CTGCTATGACCATGATTACGCCAAGCT-3′.

Substrate Complement: RNA. For assays performed using dsRNA, the substrate complement strand was obtained by ligating a chemically synthesized 20-nt RNA oligomer (Integrated DNA Technologies) carrying a 5′-biotin tag to a 187-nt RNA sequence AG(A)155 bp repeats) underscored: 5′-GTGAATTCACTCCTACAAGGAATTTGACCAAAGTATCAAAACAATTAGATCAC-3′.

The plasmid was used to generate dsRNA templates from which new substrate and substrate-complement RNAs could be transcribed. The DNA template for the substrate RNA was generated by PCR using primers 19rep-motor-NEW forward: 5′-AAAAAGAGTATAATACGACCTACTATAGGCTGACTTCTTGT-TG-3′ and 19rep-motor-NEW reverse: 5′-ATGGCATTTATATACGACCTACTATAGGCTGACTTCTTGT-TG-3′. The RNA was hybridized to a DNA complement carrying a 5′-biotin tag (Integrated DNA Technologies). The sequence was as follows (bases complementary to the substrate RNA are underscored): 5′-CCATGTTAGAAGATTCATCTTCTACACACAGA-3′.

RNase R: 1,500-bp dsRNA Constructs. Three copies of the 155-bp pALB3 sequence were inserted, in a tandem series, into a pUC19 plasmid between the EcoRI and HindIII restriction sites. The final sequence of the full 483-bp insert was as follows, with restriction sites (adjoining the 155 bp repeats) underscored: 5′-AAATCCGGCTGACTAGTCTTGGCAGAGGCGG-3′.

We note that in the dsRNA construct the sequence for the double-stranded region is frame-shifted by two base pairs relative to that of the DNA–RNA hybrid. Consequently, the substrate RNA in the former case carried a 14-nt-long 3′ overhang [sequence AG(A)12], whereas the substrate RNA in the latter case carried a 5′ GG overhang in addition to a 12-nt-long 3′ overhang (sequence A12).

DNA–RNA Hybrid: AU50 Construct. Certain experiments used a modified RNA substrate in which the first 48 bp of the DNA–RNA hybrid encountered by PNPase were replaced with a randomly generated sequence with high A+U content. The first 50 nt encountered by the ribonuclease in this construct included 35 nt that were either A or U. The sequence of this substrate (AU50) was as follows, with the randomly generated component underscored: 5′-GGCCUGACAUAGAGUUCCGGAACCGGUGGUUUGACUGGAGAAUGCAGAUAACUGGGCGGUGACCUCUUUCGGCCACCGGUGUUAAACUGCGUGUCUCUGUUAGAAUGACUGGAGUAAAGCAGGUGCCUGUAGCAUUUGCGGAGCUAGGCAGCGGAGAAGCACC-GUAAAGACAUUUAGACAGAGAAGAGAAAGA-3′.

As before, this sequence was generated by in vitro transcription of a DNA template which was obtained by PCR of pALB3 using primers RNase R forward and 48-ATrich reverse: 5′-TTTTTTTTTTTGTTAGAATTACCTTACAAGGAATTTGACTACACAGA-3′.

The DNA product (207 nt) was then used as a substrate to generate the 187-nt sequence for the RNA complement (Integrated DNA Technologies). The sequence was as follows (bases complementary to the substrate RNA are underscored): 5′-CTGCTATGACCATGATTACGCCAAGCT-3′.
RNase R data also were acquired for an alternative 1,500-bp construct carrying a single modified copy of the 155-bp pAL3 sequence in which a 25-bp segment was moved internally to a new location. To generate this modified construct, a DNA insert with the following sequence was synthesized chemically (Integrated DNA Technologies) and placed between the EcoRI and HindIII restriction sites in pUC19: 5′-ATCCCAGCTAAGCTCTCGTG-CCGCAACGGTGTGGTACGTCCGCCTGGTTAAA-CTGCGTCTGCAGGAAATGCAATTTCCGGCTATGCTATAGGCGCAGGCGAACCGGTGTTTGACGTCACCGCTGCGCGTTAAA-GCCACCTAAAAAGATCATTTAGAAAC-3′.

As above, this plasmid was used to generate DNA templates from which substrate and substrate-complement RNAs could be transcribed. The DNA template for the substrate RNA was generated by PCR using the primers pUC19-forward: 5′-AAAAAGATATAATACGACTCAC-TATAGGCTGCTCCTAATTAGCCTCGTGCCCACTGATGAGGTCTTGGTCTTGGTGTTGACGTCACCGCTGCGCGTTAAA-GCCACCTAAAAAGATCATTTAGAAAC-3′ and pUC19-reverse: 5′-GGGATTCTAAGGGAAAA-3′ overhang. For DNA substrate, its complement, and the DNA handles were annealed and ligated.

DNA Handles. The 3.057-bp and 2.250-bp digoxigenin-labeled DNA handles were derived by PCR from plasmid M13mp18 using primers L dig 3106 for the 3.057-bp handle: 5′-TTAAGGCTCTCAAC-CTCCCGCAAG-3′, L dig 4114long for the 2.250-bp handle: 5′-AAGGATTTACGGAATTAAATATACGCCGAG-3′ and P_abc_splintanneal: 5′-TCCACGATTTGCTGCTGGCCTCTG/ idSpATGTGCTGCAAGGCGATTAAGTTG-3′.

The resulting product carrying a 25 nt overhang was annealed and ligated to a DNA oligomer (Integrated DNA Technologies) of sequence 5′-AAGCGCGTACGACGATATTAGCATGCTCGA-CGGTAGAGCTTCTGCTGAGC-3′ to generate a 20 nt overhang. For DNA-RNA hybrid experiments, the D handle forward: 5′-GAGAAGGTTAATAATACGACTCAC-TATAGGCTGCTCCTAATTAGCCTCGTGCCCACTGATGAGGTCTTGGTCTTGGTGTTGACGTCACCGCTGCGCGTTAAA-GCCACCTAAAAAGATCATTTAGAAAC-3′ and D handle reverse: 5′-CACTGCGATCCTAGCCGAGGCTGTTCCGTCAGGCGACCC/ idSp/TATCCACGATTTGCTGCTGGCCTCTG/ idSp/GAGGATTTACGGAATTAAATATACGCCGAG-3′ were used to generate DNA templates from the pRL732 plasmid using primers 732-Handle forward: 5′-CACGAGAAGCTATAATACGACTCAC-TATAGGCTGCTCCTAATTAGCCTCGTGCCCACTGATGAGGTCTTGGTCTTGGTGTTGACGTCACCGCTGCGCGTTAAA-GCCACCTAAAAAGATCATTTAGAAAC-3′ and 732-Handle reverse: 5′-CTCACGAGACGCGGTTGTTCTGGG-3′.

Hybridization and Storage of Tether Components. The DNA substrate, its complement, and the DNA handles were annealed and hybridized in a solution of 80% formamide, 40 mM Pipes (pH 6.5), 400 mM NaCl, and 1 mM EDTA. Formamide was used because it reduces the thermal stability of double-stranded polynucleotides and allows annealing to be performed at lower temperatures (1). Components were brought to a temperature of 85 °C for 10 min, reduced to a temperature of 62 °C for 90 min, then to 52 °C for 90 min, and finally to 10 °C over the course of 10 min. Tethers were precipitated overnight in isopropanol at −20 °C, resuspended in TE buffer (Thermo Fischer Scientific), aliquoted, flash-frozen in liquid nitrogen, and stored at −80 °C.

Data Collection and Analysis. The optical trapping instrument has been described previously (2). Positional data were acquired at 2 kHz using custom software (LabVIEW), filtered at 1 kHz using an eight-pole low-pass Bessel filter, and boxcar averaged over a 20-point window to provide positional feedback to an active force clamp at a rate of 100 Hz. This rate is sufficient to measure speeds up to ~1,000 m/s. Uncertainties in force, caused by variations in bead size and systematic calibration errors, were estimated at 15%. Data were analyzed using Igor Pro software (WaveMetrics).

Calculation of Enzyme Velocity. Velocities of individual proteins were determined by dividing the observed change in extension by the corresponding time. Pause-free velocities for RNase R were obtained by limiting this analysis to motion over a comparatively pause-free region located 16–25 nm from the initial starting extension. This analysis was deemed preferable to one that attempted to detect and remove pauses from the individual records.

Identification of Pauses in RNase R Records. To estimate the locations of sequence-dependent pauses by RNase R on the 155-bp dsRNA construct, each record initially was truncated to include only ~1 s of positional data before the start of processive unwinding and ~0.5 s after the cessation of enzyme motion. A histogram of occupancy versus position (bin size, 0.1 nm) was constructed for each trace, with peaks corresponding to enzyme pauses. The baseline starting position for each record was identified by fitting a Gaussian function to the first peak present (corresponding to the initial ~1 s before motion) to allow alignment of traces. Once aligned, the data were summed for all records, with the contribution of each trace weighted by its integrated occupancy over the entire record (excluding the contribution of the initial and final peaks) to minimize the influence of outliers. The resulting positional occupancy histogram displayed three well-defined peaks, corresponding to pause positions for RNase R, the centers of which were identified by Gaussian fits. This analysis returned peaks at the same positions when performed using an unsupervised weight of the individual records, and the calculation was robust against minor changes in histogram bin size.

A similar calculation was performed for records of RNase R over the 1,500-bp dsRNA construct carrying tandem repeats of the 155-bp sequence. Here, the individual contributions of records were weighted by the integrated occupancy over the first 20 nm traversed because not all enzymes moved beyond this location. To adjust for possible bias in the positional occupancy arising from only a subset of motors advancing to a given position, the occupancy at each position in the cumulative histogram was normalized relative to the number of enzymes that unwound the RNA up to that point.

Periodogram Analyses. To investigate stepping behavior, records were truncated to include only 0.5 s of positional data before processive motion, and the terminal backtracks of 12 ± 1 nt observed in records of PNPase (see main text) were excluded from the analysis. We calculated the positional separations of all points relative to each other within each record (i.e., pairwise distances), and histogrammed these positions (bin size, 0.001 nm) to determine the distribution of pairwise positions. These data then were summed across equivalent single-molecule records, with each record weighted by its respective number of pairwise separations, to ensure that each record contributed appropriately to the analysis and to minimize the influence of any potential outliers. A Fourier transform of this distribution was performed to identify peaks in spatial frequency. Each peak was fit to a Gaussian function to identify its center, and the SD of a peak was taken to represent an estimate of the positional error. Given the levels of signal and noise, we estimate that our system would not detect steps smaller than three nucleotides.

Pause locations for RNase R on the 155-bp construct. Filled boxes (red) above the sequence indicate the locations of the three pauses (pause 1, pause II, and pause III) along with positional uncertainties (error bars). Two nucleotides at the 5′ end and the 12 nt at the 3′ end (colored blue) represent overhang sequences that do not hybridize with the complementary strand.

Single-molecule assay—the 1,500-bp dsRNA construct. As in Fig. 1 in the main text, a double-stranded segment of RNA is placed under tension between two polystyrene beads (light blue) that are held in separate optical traps (pink). One strand of RNA (the digested strand, orange) carries a 3′, adenine-rich terminal overhang that serves as a loading site for RNase R. The complementary strand (red) is hybridized at both ends to DNA handles that are coupled to separate beads via biotin–avidin and digoxygenin–antibody linkages. The RNA immediately adjacent to the 3′ overhang consists of a 26-bp spacer element followed by three copies of the 155-bp sequence used previously. Tandem repeats (indicated) are separated by six-base pair restriction sites (red).

Periodogram analysis of records of RNase R movement (green curve; n = 13). Analysis was performed on data collected from the 155-bp dsRNA construct. Unlike the corresponding periodograms for PNPase, no strong peak is evident at a position indicating steps of six or seven nucleotides (vertical gray dashed line, positioned as in Fig. 3B in the main text). The two peaks in the distribution at 0.16 nm$^{-1}$ and 0.31 nm$^{-1}$ correspond to the larger ∼6-nm and ∼3-nm separations found between sequence-dependent pauses, as reported earlier (Fig. 2B in the main text).

Five representative single-molecule records of mutant PNPase (ΔKH–ΔS1) activity on 155-bp dsRNA constructs. Enzymes moved an average distance of 14 ± 1 bp (mean ± SE). Individual traces are displayed in different colors and are offset horizontally.